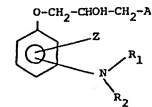
UK Patent Application (19) GB (11) 2 129 022 A

- (21) Application No 8331092
- (22) Date of filing 25 Jan 1983 Date lodged 22 Nov 1983
- (30) Priority data
- (31) 83900
- (32) 26 Jan 1982
- (31) 84391
- (32) 27 Sep 1982
- (33) Luxembourg (LU)
- (43) Application published 10 May 1984
- (51) INT CL³ A61K 7/13
- (52) Domestic classification D1B 2A4 2D
- (56) Documents cited None
- (58) Field of search D1B
- (60) Derived from Application No 8301981 under Section 15(4) of the Patents Act 1977
- (71) Applicant
 L'Oreal,
 (France),
 14 rue Royale,
 75008 Paris,
 France
- (72) Inventors
 Andree Bugaut,
 Alain Genet
- (74) Agent and/or address for service
 J. A. Kemp & Co.,
 14 South Square,
 Gray's Inn,
 London,
 WC1R 5EU

(54) Hair dyeing compositions containing 1-(substituted phenoxy)-'3-amino-propan-2-ol compounds

(57) A hair dyeing composition comprises at least one compound or salt thereof of formula



in which

Z represents NO,

Z may also represent an amino group depending on certain provisos, A represents NY or



wherein Y represents two identical or different C_{1-4} alkyl or C_{1-4} hydroxyalkyl groups or together with the nitrogen of NY or



represent a morpholino or piperidino ring or, when A represents the group NY, may also represent two hydrogen atoms or a hydrogen atom and a C_{1-4} hydroxyalkyl group, Y' represents a C_{1-4} alkyl group and X represents an anion, and R_1 and R_2 each independently represents H, C_{1-4} alkyl or C_{1-4} hydroxyalkyl, together with an appropriate carrier.

10

15

20.

25

30

SPECIFICATION

New compounds which can b us d for hair dy ing, proc ss for th ir pr paration, dyeing compositions in which they are present and corresponding hair-dyeing process

The present invention relates to new 1-(substituted phenexy)-3-aminopropan-2-ol compounds in 5 which the extra-nuclear amine group may or may not be substituted, and to the process for their preparation. The invention also relates to hair-dyeing compositions containing these new compounds and to a dyeing process using the said compositions.

The object of the invention is to propose a new class of compounds which can be used in hair dyes to give strong colourations which are stable to light and have a good fastness to weather and washing. A further object of the invention is to propose a new category of compounds having a good degree of harmlessness and satisfactory characteristics from the point of view of mutagenesis.

The present invention relates to a new chemical compound of the formula (I) (or a corresponding acid salt)

in which formula:

Z represents NO2 or NH2;

A denotes NY or the group

Y representing two identical or different lower alkyl or lower hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a morpholine or piperidine 20 heterocyclic ring with the nitrogen atom to which they are attached, and it also being possible for one or both of the substituents of which Y is composed to represent hydrogen, only if A denotes NY, Y' representing a lower alkyl substituent group having at most 4 carbon atoms, and X representing an anion; and

— R₁ and R₂ are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon atoms, with the proviso that if A denotes NY, 1°) if Z is in the 4position on the benzene nucleus and if NR₁R₂ is in the 2-position, R₁ and R₂ both representing a hydrogen atom, one of the two substituents of which Y is composed does not represent an ethyl group if the other substituent represents a hydrogen atom, and the two substituents of which Y is composed do not form a morpholine or piperidine heterocyclic ring with the nitrogen atom to 30 which they are attached, and 2°) if one of the two substituents of which Y is composed denotes a

hydrogen atom or an alkyl group and the other denotes an alkyl group, Z cannot be located in the 3-position and NR₁R₂ in the 2-position R₁ denoting a hydrogen atom and R₂ denoting a hydrogen atom or an alkyl group.

35 X can advantageously be a halogen, in particular chlorine. 35 The present invention also relates to a process for the preparation of a chemical compound of the formula (I) (or a corresponding acid salt) in which:

Z represents NO2 or NH2;

45

A denotes NY, Y representing two identical or different lower alkyl or lower hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a 40 40 morpholine or piperidine heterocyclic ring with the nitrogen atom to which they are attached, and it also being possible for one or both of the substituents of which Y is composed to represent hydrogen; and

 R_1 and R_2 represent hydrogen atoms, with the proviso that if Z is in the 4-position on the benzene nucleus and if NR_1R_2 is in the 2-position, one of the two substituents of which Y is composed does not represent an ethyl group if the other substituent represents a hydrogen atom, and the two substituents of which Y is composed do not form a morpholine or piperidin ring with the nitrogen atom to which they are attached; the process comprising the following steps:

10

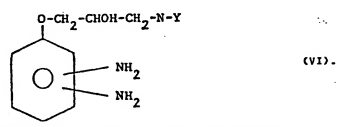
a) epichlorohydrin is react d with the known compound of the formula (II)

to give the compound of the formula (III)

b) an amine of the formula HNY, in which Y has the meaning indicated above, is reacted with the compound of the formula (III) to give the compound of the formula (IV)

c) a strong acid, such as hydrochloric acid, is reacted with the compound of the formula (IV) to give the compound of the formula (V)

d) and, if appropriate, to obtain the compound of the formula (I) in which Z represents NH₂, reduction is carried out either, for example, by means of zinc powder in an alcoholic medium in the presence of ammonium chloride, or by means of catalytic hydrogenation, it being possible for palladium-on-charcoal to be used as the catalyst, and this makes it possible to obtain the compound of the formula (VI)



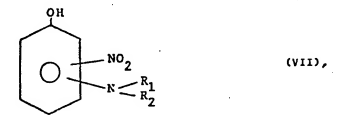
The pr sent invention also relates to a proc ss for the pr paration of a chemical compound f the formula (I) (or c rresponding acid salt) in which:

Z represents NO₂ or NH₂;

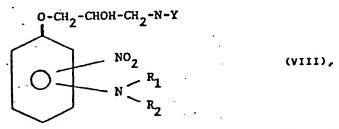
25

— A den tes NY, Y representing twildentical r different lower alkyl or low r hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a morpholine or piporidine heterocyclic ring with thousand it also being possible for one or both of the substituents of which Y is composed to represent hydrogen; and

R₁ and R₂ are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon atoms, the case in which R₁=R₂=H being excluded, in which process the compound of the formula (VII)



in which formula R₁ and R₂ have the meanings indicated above, is used as the starting material, and the reactions involved in steps a) and b) mentioned above are carried out successively to give a product of the formula (VIII)



in which Y, R₁ and R₂ have the meanings indicated above, and if it is desired to obtain a compound of the formula (I) (or a corresponding acid salt) in which Z represents NH₂, A denoting NY, the compound of the formula (VIII) is subjected to reduction as indicated in step d) mentioned above. As a variant, if neither of the substituents of which Y Is composed represents a hydrogen atom, it is possible to use a process of preparation which starts from a compound of the formula (II), to carry out steps a), b), c) and d) and then to introduce a substituent into one of the nuclear amine groups of the compound of the formula (VI), for example by introducing a substituent into the corresponding arylsulphonamide and subjecting this substituted aryl sulphonamide to acid hydrolysis.

The present invention also relates to a process for the preparation of a compound of the formula (I) in which:

25 — Z denotes NO₂ or NH₂;

- A denotes

30

35



Y representing two identical or different lower alkyl or lower hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a morpholine or piperidine heterocyclic ring with the nitrogen atom to which they are attached, Y' representing a lower alkyl substituent group having at most 4 carbon atoms, and X representing an anion; and R₁ and R₂ are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon at ms; in which process the nuclear amine group (or groups) of the corresponding tertiary compound is (or are) protected by acetylation, this being followed by reaction with a quat rnising agent. Subsequently, the product is deacetylated by reaction with an acid and, if appropriate, the quaternisation anion can be changed at the same time.

In the case wher Z repr sents a nitro group, the compounds of the formula (I) are direct dyestuffs which can be used for dyeing keratin fibres, and in particular hair. It has been found that

th se compounds have the advantage of good solubility in the solv into generally used in hair dyeing, and this enables them to be used at a sufficiently high concentration to give the keratin fibre a strong colouration with a good uniformity. The dyeing obtained has a good stability to light and weather. Furthermore, the use of these compounds in hair dyeing is characterised by a good degree of harmlessness. The invention thus also relates to a dyeing composition for keratin fibres, and in particular for hair, which composition contains, in an appropriate carrier, at least one compound of the formula (I) (or a corresponding acid salt) in which:

5

- Z represents a nitro group;

— A denotes NY or

10

15

20



10

Y representing two identical or different lower alkyl or lower hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a morpholine or piperidine heterocyclic ring with the nitrogen atom to which they are attached, and it also being possible for one or both of the substituents of which Y is composed to represent hydrogen, only if A denotes NY, Y' representing a lower alkyl substituent group having at most 4 carbon atoms, and X representing an anion; and

15

— R₁ and R₂ are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon atoms. In a preferred embodiment, the dyeing composition according to the invention contains from 0.001% to 4% by weight of at least one compound of the formula (I).

20

In the case where Z represents an NH₂ group, the compounds of the formula (I) are either oxidation bases, in the case of para-diamines or ortho-diamines, or couplers intended to be used in association with oxidation bases, in the case of meta-diamines.

If the compound of the formula (I) is an oxidation base, the invention also relates to a dyeing composition for keratin fibres, in particular for hair, which composition contains, in an appropriate carrier, at least one compound of the formula (I) (or a corresponding acid salt), in which formula:

25

- Z represents an NH₂ group;

- A denotes NY or

Y representing two identical or different lower alkyl or lower hydroxyalkyl substituent groups having at most 4 carbon atoms, these two groups optionally forming a morpholine or piperidine heterocyclic ring with the nitrogen atom to which they are attached, and it also being possible for one or both of the substituents of which Y is composed to represent hydrogen, only if A denotes NY, Y' representing a lower alkyl substituent group having at most 4 carbon atoms, and X representing an anion; and

30

representing an anion; and

R₁ and R₂ are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon atoms. In a preferred embodiment, the compound of the formula (I) is used in a proportion of between 0.001% and 4% by weight, relative to the total weight of the composition.

35

In the case where the compound of the formula (I) is a meta-phenylenediamine, the invention
also relates to a dyeing composition for keratin fibres, and in particular for hair, containing, in an
appropriate carrier, at least one oxidation base, which composition also contains at least one
compound of the formula (I) (or a corresponding acid salt), in which formula:

40

Z repr sents an NH₂ group;

45 - A denotes NY or



10

15

20

25

35

5

10

30

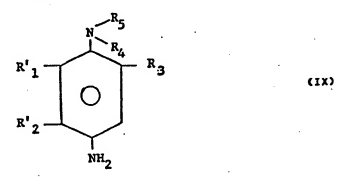
35

Y representing two identical or different lower alkyl or lower hydr xyalkyl substituent groups having at most 4 carbon atoms, these two gr ups opti nally forming a morpholine or piperidine h terocyclic ring with the nitrogen atom to which they are attached, and it also being possible for ne or both of the substituents of which Y is composed to r present hydrogen, only if A den tes NY, Y' representing a lower alkyl substituent group having at most 4 carbon atoms, and X representing an anion; and

representing an anion; and R_1 and R_2 are identical or different and represent a hydrogen atom or a lower alkyl or hydroxyalkyl radical having at most 4 carbon atoms, with the proviso that if A denotes NY, if Z is in the 4-position on the benzene nucleus and if NR_1R_2 is in the 2-position, R_1 and R_2 both representing a hydrogen atom, one of the two substituents of which Y is composed does not represent an ethyl group if the other substituent represents a hydrogen atom, and the two substituents of which Y is composed do not form a morpholine or piperidine heterocyclic ring with the nitrogen atom to which they are attached. The compound of the formula (I) is preferably used in a proportion of between 0.001% and 2.5% by weight, relative to the total weight of the composition.

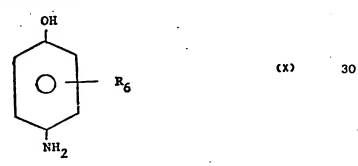
The dyeing compositions according to the invention can comprise, in addition to the compounds of the formula (I), oxidation bases consisting of:

A-Para-phenylenediamines of the general formula (IX)



or the corresponding acid salts, in which formula R'₁, R'₂ and R₃ are identical or different and represent a hydrogen atom, an alkyl radical having from 1 to 4 carbon atoms, an alkoxy radical having 1 or 2 carbon atoms or a halogen atom, and R₄ and R₅ are identical or different and represent a hydrogen atom, an alkyl or hydroxyalkyl radical, an alkoxyalkyl radical in which the alkoxy group contains 1 or 2 carbon atoms, or a carbamylalkyl, alkylsulphonamidoalkyl, acetylaminoalkyl, ureidoalkyl, carbethoxyaminoalkyl, aminoalkyl, monoalkylaminoalkyl, dialkylaminoalkyl, piperidinoalkyl or morpholinoalkyl radical, the alkyl groups in R₄ and R₅ having from 1 to 4 carbon atoms, or alternatively R₄ and R₅ form a piperidino or morpholino group together with the nitrogen atom to which they are attached, with the proviso that R'₁ and R₃ represent a hydrogen atom if R₄ and R₅ do not represent a hydrogen atom.

B — Para-aminophenols of the general formula (X)



or the corresponding acid salts, in which formula R_6 represents a hydrogen atom, an alkyl radical containing from 1 to 4 carbon atoms or a halogen atom such as, for example, chlorine or bromine.

C — Heterocyclic bases such as 2,5-diaminopyridine, 3-m thyl-7aminobenzomorpholine and 5-aminoindole.

In additi n to the compounds f th formula (I), the dy ing compositions according to the invention can contain the following products, taken in isolation r in combination:

1) if the composition contains at least ne xidation bas : at least one c upler taken from the group comprising resorcinol, pyrocatechol, 2-m thylresorcin l, 2-ethylresorcinol, meta-aminophenol, 2-m thyl-5-aminophenol, 2-methyl-5-N- $(\beta$ -hydroxyethyl)-amin phenol, 6-hydroxybenzomorpholin ,

GB 2 129 022 A 6

2,6-dimethyl-3-acetylaminophenol, 2-m thyl-5-carbethoxyaminophenol, 2-methoxy-5-carbethoxyaminoph nol, 2-methyl-5-ureidophenol, 2,4-diaminoph noxy thanol, 2,4-diaminoanisole, 2,6dimethyl-meta-ph nylen diamine, 2-amino-4-N-methylamin phenoxyethanol, 2,4-diaminoph nyl β methoxyethyl eth r, 2,4-diaminophenyl \(\beta\)-mesylaminoethyl eth r, 2-N-carbamylmethylamino-4-5 aminoanisole, 3-amin -4-methoxyphenol, α -naphthol, 2,6-diaminopyridine, 3,5-diamino-2,6dimethoxypyridine, 1-phenyl-3-methylpyrazol-5-one and 2-amino-4-N-(β -hydroxyethyl)-aminophenyl β-hydroxypropyl ether; 2) ortho-phenylenediamines and ortho-aminophenols optionally containing substituents on the nucleus or on the amine groups, or ortho-diphenol, it being possible, by means of complex oxidation 10 mechanisms, for these products to lead to new coloured compounds, either by cyclisation with themselves or by reaction with para-phenylenediamines; 3) dyestuff precursors of the benzene series, containing, on the nucleus, at least three substituents chosen from the group comprising hydroxyl, methoxy or amino groups, such as 2,6diaminohydroquinone dihydrochloride, 2,6-diamino-4-N,N-bis-(ethyl)-aminophenyl trihydrochloride, 15 2,4-diaminophenol dihydrochloride, 1,2,4-trihydroxybenzene, 2,3,5-trihydroxytoluene or 4-methoxy-2-15 amino-N-(β-hydroxyethyl)-aniline; 4) quinone dyestuffs such as 2-hydroxy-1,4-naphthoquinone, 5-hydroxy-1,4-naphthoquinone and 2-[4'-(N,N-dihydroxyethylamino)-anilino]-5-N'-(β-hydroxyethyl)-amino-1,4-benzoquinone; 5) indoanilines, indophenols or indamines, or their leuco derivatives, such as 4,4'-dihydroxy-2-20 20 amino-5-methyldiphenylamine, 4,4'-dihydroxy-2-N-(β-hydroxyethyl)-amino-5-methyl-2'-chlorodiphenylamine, 2,4'-diamino-4-hydroxy-5-methyldiphenylamine, 2,4-dihydroxy-4'-N-(βmethoxyethyl)-aminodiphenylamine and 2,4-dihydroxy-5-methyl-4'-N-(β-methoxyethyl)-aminodiphenylamine; 6) direct dyestuffs chosen from the class of the azo dyestuffs, the anthraquinone dyestuffs and 25 the nitro dyestuffs of the benzene series, such as 2-amino-3-nitrophenol, 1-amino-2-nitro-4N-(β-25 hydroxyethyl)-amino-5-methylbenzene, 1-N,N-bis-(eta-hydroxyethyl)-amino-3-nitro-4-N'-methylaminobenzene, 1-N-methyl-N-(β-hydroxyethyl)-amino-3-nitro-4-N'-(β-hydroxyethyl)-aminobenzene, 1-Nmethyl-N-(β-hydroxyethyl)-amino-3-nitro-4-N'-methylaminobenzene, 3-nitro-4-N-(β-hydroxyethyl)aminoanisole. 3-nitro-4-N-(β-hydroxyethyl)-aminophenol, 3-nitro-4-aminophenoxyethanol, 3-nitro-4-30 N-methylaminophenoxyethanol, 3-N-methylamino-4-nitrophenoxyethanol, 3-nitro-4-N-(βaminoethyl)-aminophenoxyethanol, 2-N-(β-hydroxyethyl)-amino-5-nitroanisole and 1,4,5,8-tetraaminoanthraquinone; and 7) various customary adjuvants such as water, penetrating agents, surface-active agents, thickeners, anti-oxidants, alkalising or acidifying agents, perfumes, sequestering agents, film-forming 35 35 products and treating agents. The pH of the dyeing compositions according to the invention is between 5 and 11.5. Amongst the alkalising agents which can be used, there may be mentioned ammonia, alkylamines such as ethylamine or triethylamine, alkanolamines such as mono-, di- or tri-ethanolamine, alkylalkanolamines such as methyldiethanolamine, the hydroxides of sodium or potassium and the carbonates of sodium, 40 potassium or ammonium. Amongst the acidifying agents which can be used, there may be mentioned 40 lactic acid, acetic acid, tartaric acid and phosphoric acid. Water-soluble anionic, cationic, non-ionic or amphoteric surface-active agents, or a mixture thereof, can also be added to the composition according to the invention. Amongst the surface-active agents which can be used in particular, there may be mentioned alkylbenzenesulphonates, alkylnaphthalenesulphonates, fatty alcohol sulphates, ether-sulphates and sulphonates, quaternary 45 ammonium salts such as trimethylcetylammonium bromide and cetylpyridinium bromide, fatty acid diethanolamines or polyoxyethyleneated or polyglycerolated acids, alcohols or alkylphenols. Preferably, the surface-active agents are present in the composition according to the invention in a proportion of between 0.5 and 55% by weight and advantageously of between 4 and 40% by weight, relative to the 50 50 total weight of the composition. Organic solvents can also be added to the composition according to the invention, and examples of these which may advantageously be mentioned are ethanol, isopropanol, glycerol, glycols and their ethers, such as 2-butoxyethanol, ethylene glycol, propylene glycol, and diethylene glycol monoethyl ether and monomethyl ether, and similar solvents. The solvents can advantageously be present in the composition in a proportion ranging from 1 to 40% by weight and preferably of between 5 and 30% by weight, relative to the total weight of the composition. The thickening products which can be added to the composition according to the invention can advantageously be taken from the group comprising sodium alginate, gum arabic, cellulose derivatives such as m thylcellulose, hydroxyethylc llulose, hydroxypropylmethylcellulose and the sodium salt of 60 carb xymethylcellul se, and acrylic acid polymers; it is also possible to use inorganic thickeners such 60 as bentonite. Preferably, the thick ners are pres int in a proportion of between 0.5 and 5% by weight, relative t the total weight of the composition, and advantageously of between 0.5 and 3% by weight. The antifixidants which can be added to the composition according to the invention can advantageously be tak in from the group comprising sodium sulphite, thioglyc lic acid, 65 mercaptosuccinic acid, sodium bisulphite, ascorbic acid and hydroquinon . Thes antioxidants can be 65

10

15

present in the composition in a proportion of between 0.05 and 1.5% by weight, relative to the total weight of the composition.

If the dyeing composition according t the invention c ntains at least one oxidation base, it contains, at the time of use, oxidising agents such as hydrogen peroxide, urea per xide or per-salts such as amm nium persulphat.

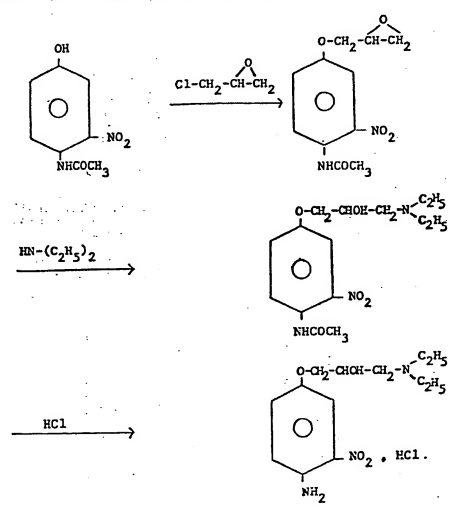
The dyeing composition according to the invention can be present in the form of a liquid, a cream, a gel or an aerosol or in any other form suitable for dyeing keratin fibres.

The present invention also relates to a new hair-dyeing process, characterised in that the dyeing composition defined above is left to act on the hair for an application time varying between 10 and 45 minutes, and in that the hair is rinsed, optionally washed and rinsed again, and dried.

In the case where the dyeing composition used contains at least one oxidation base, the abovementioned process includes an initial stage in which a sufficient amount of oxidising agents is mixed with the said composition at the time of use.

To provide a clearer understanding of the object of the invention, several embodiments thereof will now be described by way of purely illustrative and non-limiting examples.

Example 1
Preparation of 1-(3'-nitro-4'-aminophenoxy)-3-N,N-diethylaminopropan-2-ol



First st p
20 Preparation of 1-(3'-nitro-4'-acetylaminophenoxy)-2,3-epoxypropane

150 ml of epichl rohydrin are added, at ambient temperature, to a solution of 0.376 mol (73.7 g) of 3-nitro-4-acetylaminophenol in 375 ml of 1.1 N sodium hydroxide solution. The reaction medium is left to stand at ambient temp rature for 48 hours, with thorough stirring, and the expected product which has precipitated is the n filtered off. After drying in vacuo and recrystallisation from benzene, this product melts at 123°C.

20

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₁ H ₁₂ N ₂ O ₅	Found
C%	52.38	52.32
H%	4.80	4.77
N%	11.11	11.06
0%	31.72	31.57

Second step

5

10

Preparation of 1-(3'-nitro-4'-acetylaminophenoxy)-3-N,N-diethylaminopropan-2-ol

0.2 mol (50.4 g) of 1-(3'-nitro-4'-acetylaminophenoxy)-2,3-epoxypropane is heated for 3 hours under reflux in 146 g of diethylamine and 50 ml of absolute alcohol. The diethylamine and the alcohol are driven off in vacuo. The residual oil crystallises slowly. After recrystallisation from cyclohexane, 58 g of the expected product, which melts at 77°C, are obtained.

The product obtained corresponds to the following analysis results:

	Analysis	Calculated for C ₁₅ H ₂₃ N ₃ O ₅	Found
ĺ	C%	55.37	55.40
١.	Н%	7.13	7.18
ı	N%	12.92	12.86
1	0%	24.59	24.78
1		1	

Third step
Preparation of 1-(3'-nitro-4'-aminophenoxy)-3-N,N-diethylaminopropan-2-ol hydrochloride
0.152 mol (49.7 g) of 1-(3'-nitro-4'-acetylaminophenoxy)-3-N,N-diethylaminopropan-2-ol is
heated for 30 minutes, in a boiling water-bath, in 100 ml of hydrochloric acid (specific gravity=1.18).
After the addition of ammonia to the reaction medium so as to give a pH of 6, the 1-(3'-nitro-4'aminophenoxy)-3-N,N-diethylaminopropan-2-ol monohydrochloride crystallises. It is filtered off,
washed with a small amount of 80° strength alcohol, recrystallised from a mixture of water and
ethanol and dried in vacuo. It melts at 178°C.

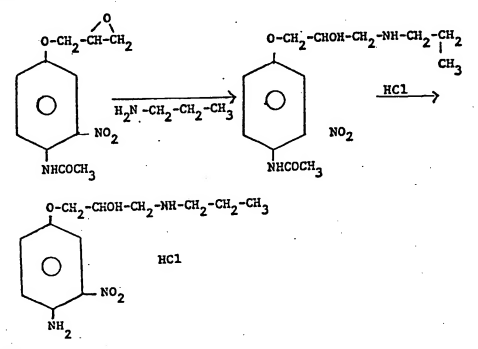
The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₃ H ₂₁ N ₃ O ₄ . HCl	Found
С%	48.82	48.65
Н%	6.93	6.90
N%	13.14	12.90
0%	20.01	20.17
CI%	11.09	10.96

20

15

Example 2
Preparation of 1-(3'-nitro-4'-aminophenoxy)-3-N-propylaminopropan-2-ol monohydrochloride



First step

5 Preparation of 1-(3'-nitro-4'-acetylaminophenoxy)-3-N-propylaminopropan-2-ol
0.02 mol (5.0 g) of 1-(3'-nitro-4'-acetylaminophenoxy)-2,3-epoxypropane, obtained according to
the first step of Example 1, is introduced into 25 ml of N-propylamine and the mixture is then heated
for 3 hours at 45°C. The reaction medium is subsequently poured into 100 g of iced water, and the
expected product which has precipitated is then filtered off and washed with water. After drying in
vacuo and recrystallisation from a mixture of benzene and ethyl acetate, the product melts at 146°C.
The product obtained corresponds to the following analysis results:

Analysis	Calculated for $C_{14}H_{21}N_3O_5$	Found
C%	54.01	54.00
H%	6.80	6.86
N%	13.50	13.36
0%	25.70	25.88
	l .	i .

Second step

15

20

Preparation of 1-(3'-nitro-4'-aminophenoxy)-3-N-propylaminopropan-2-ol monohydrochloride 0.008 mol (2.5 g) of 1-(3'-nitro-4'-acetylaminophenoxy)-3-N-propylaminopropan-2-ol is heated for 30 minutes, in a boiling water-bath, in 7 ml of hydrochloric acid (specific gravity=1.18), with stirring. The cooled reaction mixture is treated with 22°Be ammonia solution up to a pH of 5. the monohydrochloride of the expected product precipitates. It is filtered off, washed with a small amount of 80° strength alcohol and then recrystallised from a mixture of water and ethanol. It melts at 224°C. The product obtained corresponds to the following analysis results:

20

15

5

Analysis	Calculated for C ₁₂ H ₁₉ N ₃ O ₄ . HCl	Found
C%	47.14	47.24
H%	6.59	6.57
N%	13.74	13.75
O%	20.93	20.76
CI%	11.60	11.55

Example 3
Pr parati n f1-(3'-nitro-4'-aminophen xy)-3-aminopropan-2-ol

First step

5 Preparation of 1-(3'-nitro-4'-acetylaminophenoxy)-3-succinimido-3-propan-2-ol

50.4 g (0.2 mol) of 1-(3'-nitro-4'-acetylaminophenoxy)-2,3-epoxypropane (prepared according to the first step of Example 1) are introduced into 150 ml of absolute ethanol to which 12 drops of pyridine have been added. 0.24 mol (23.7 g) of succinimide is added. The mixture is heated under reflux for 5 hours 30 minutes. On cooling, the expected product crystallises. It is filtered off and washed with a small amount of alcohol. After recrystallisation from alcohol and drying in vacuo, it melts at 154°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₅ H ₁₇ N ₃ O ₇	Found
• С%	51.28	51.34
Н%	4.88	4.95
N%	11.96	11.86
. 0%	31.83	31.84

Second step

15 Preparation of 1-(3'-nitro-4'-aminophenoxy)-3-aminopropan-2-ol

50.9 g (0.145 mol) of the succinimido derivative obtained according to the first step are introduced into 100 ml of 96° strength alcohol to which 250 ml of hydrochloric acid (specific gravity=1.18) have been added. After 14 hours under reflux, the cool d reaction medium is treated with acetone. The expected product precipitates in the form of the hydrochloride. This hydrochloride is taken up in 70 ml of water. By rendering the aque us solution alkaline with 10 N sodium hydroxide s lution, the 1-(3'-nitro-4'-aminoph noxy)-3-aminopropan-2-ol is precipitated in the form of an oil, which crystallis s rapidly. After r crystallisation from alcohol, the product m Its at 124°C.

15

5

10

The product obtain d corresponds to the following analysis results:

	Analysis	Calculated for C ₈ H ₁₃ N ₃ O ₄	Found
I	C%	47.57	47.54
ı	Н%	5.77	5.73
ı	N%	18.49	18.36
ı	0%	28.17	28.18
1			

Example 4 Preparation of 1-(2'-amino-4'-nitrophenoxy)-3-aminopropan-2-ol

First step

Preparation of 1-(2'-acetylamin -4'-nitroph noxy)-2,3-epoxypropane

1.6 litres of epichlorohydrin are added to a solution of 2 mols (392 g) of 2-acetylamin -4nitrophenol in two litres of 1.1 N sodium hydroxide solution, and the reaction medium is left to stand for 96 h urs at 20°C, with thorough stirring. 282 g of the expected product which has precipitated are filtered off and washed with water. This product contains about 20% of a by-product, namely 1,3-bis(2'-acetylamino-4'-nitrophenoxy)-propan-2-ol (m lting at 233°C), which results from the condensation of one molecul of pichlorohydrin with two mols of 2-acetylamino-4-nitrophenol. H wever, it will be used as such for the second step of the synthesis, the by-product being very readily removed at this stag.

5 Second step

5

Preparation of 1-(2'-acetylamino-4'-nitrophenoxy)-3-succinimidopropan-2-ol

100 g of the crude product obtained in the previous step are introduced into 500 ml of 95° strength ethanol. 0.52 mol (51.5 g) of succinimide and 2.5 ml of pyridine are added and the mixture is heated under reflux for 2 hours, with stirring. The alcoholic solution is filtered at the boil to isolate the expected product, which is insoluble in hot alcohol. This gives 60 g of virtually pure 1-(2'-acetylamino-4'-nitrophenoxy)-3-succinimidopropan-2-ol. After recrystallisation from alcohol, this product melts at 184°C.

10

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₅ H ₁₇ N ₃ O ₇	Found
С%	51.28	51.15
Н%	4.84	4.87
N%	11.96	11.86
0%	31.91	32.15
	1	i .

The mother liquors from the reaction medium contain a small amount of succinimido derivative together with 1.3-bis-(2'-acetylamino-4'-nitrophenoxy)-propan-2-ol, which has thus been removed.

15

Third step

Preparation of 1-(2'-amino-4'-nitrophenoxy)-3-succinimidopropan-2-ol

2 g (0.0057 mol) of the 1-(2'-acetylamino-4'-nitrophenoxy)-3-succinimidopropan-2-ol obtained in the second step are heated for one hour, in a boiling water-bath, in 10 ml of hydrochloric acid (specific gravity=1.18), with stirring. After the reaction medium has been cooled, diluted and rendered alkaline, the expected product precipitates. It is filtered off, washed with water and recrystallised from ethanol. It melts at 200°C.

20

The product obtained corresponds to the following analysis results:

25

Analysis	Calculated for C ₁₃ H ₁₅ N ₃ O ₆	Found
С%	50.48	50.47
H%	4.85	4.87
N%	13.59	13.58
0%	31.07	30.94

25

Fourth step

Preparation of 1-(2'-amino-4'-nitrophenoxy)-3-aminopropan-2-ol

7.9 g (0.0256 mol) of the succinimido derivative prepared in the previous step are heated for 8 hours in 40 ml of hydrochloric acid (specific gravity=1.18) under reflux, with stirring. After the reaction medium has cooled, the expected product, which has crystallised in the form of the dihydrochloride, is filtered off. This dihydrochloride is dissolved in 175 ml of water and the pH is brought to 9 with sodium hydroxide solution. The expected product which has precipitated is filtered off. After washing with water and recrystallisation from alcohol, the product melts at 159°C.

30

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₉ H ₁₃ N ₃ O ₄	Found
· C%	47.58	47.57
· H%	5.73	5.75
N%	18.50	18.56
0%	28.19	28.16

35

Example 5 Preparation of 1-(2'-amino-5'-nitroph n xy)-3-aminopropan-2-ol

First step

Preparation of 1-(2'-acetylamino-5'-nitrophenoxy)-2,3-epoxypropane 0.376 mol (73.7 g) of 2-acetylamino-5-nitrophenol is dissolved in 375 ml of 1.1 N sodium hydroxide solution, and 300 ml of epichlorohydrin are then added. The reaction medium is left to stand for 96 hours at 20°C, with thorough stirring, and the expected product which has precipitated is then filtered off and washed with water. After recrystallisation from ethanol and drying in vacuo, it melts at 10 164°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₁ H ₁₂ N ₂ O ₅	Found
C%	52.38	52.29
H%	4.80	4.83
N%	11.11	11.20
O%	31.72	31.86

Second step

15

Pr paration f1-(2'-acetylamino-5'-nitrophenoxy)-3-succinimidopropan-2-ol 0.072 mol (18.2 g) of 1-(2'-acetylamino-5'-nitrophenoxy)-2,3-epoxypropane is introduced into

60 ml f absolut ethanol to which 6 drops of pyridin have been added. 0.087 mol (8.6 g) of succinimide is added and the mixture is then heated undir reflux for 5 hours. After cooling, 150 ml of water are added and the alcohol is driven off in vacuo. The expected product, which is obtained initially in the form of a water-insoluble oil, crystallises slowly. It is filtered off and recrystallis d from 96° 20 strength alcohol. After drying, it melts at 157°C.

15

5

10

15

Analysis	Calculated for C ₁₅ H ₁₇ N ₃ O ₇	Found
C%	51.28	51.30
Н%	4.88	4.94
N%	11.96	11.89
0%	31.88	32.07

Third step

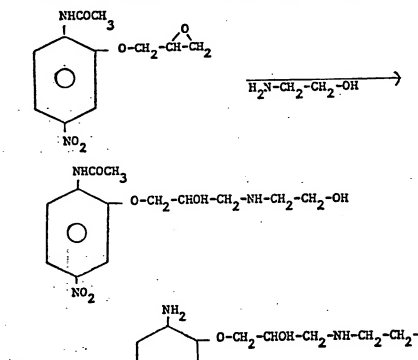
Preparation of 1-(2'-amino-5'-nitrophenoxy)-3-aminopropan-2-ol

7.0 g (0.02 mol) of the succinimido derivative obtained in the second step are heated under reflux for 14 hours in 14 ml of 96° strength alcohol to which 38 ml of hydrochloric acid (specific gravity=1.18) have been added. The reaction medium is then cooled and treated with acetone in order to precipitate the expected product in the form of the hydrochloride. This hydrochloride is filtered off and then dissolved in 12 ml of water. By rendering the aqueous solution alkaline with 10 N sodium hydroxide solution, the 1-(2'-amino-5'-nitrophenoxy)-3-aminopropan-2-ol is precipitated in the form of an oil, which crystallises rapidly. The product is filtered off, washed with water and recrystallised from ethanol. It melts at 141°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₉ H ₁₃ N ₃ O ₄	Found
С%	47.57	47.58
Н%	5.77	5.76
N%	18.49	18.62
0%	28.17	28.28

15 Example 6 Preparation of 1-(2'-amino-5'-nitrophenoxy)-3-N-(β-hydroxyethyl)-aminopropan-2-ol



$$\begin{array}{c}
 & \text{NH}_2 \\
 & \text{O-CH}_2\text{-CHOH-CH}_2\text{-NH-CH}_2\text{-CH}_2\text{-OH} \\
 & \text{NO}_2
\end{array}$$

20

Preparation of 1- (2'-acetylamino-5'-nitrophenoxy)-3-N-(β-hydroxyethyl)-aminopropan-2-ol

10.1 g (0.04 mol) of the 1-(2'-acetylamin -5'-nitrophenoxy)-2,3-epoxypropane prepared according to the first stage of Example 5 are dissolv d in 36.6 g of monoethanolamine and 10 ml f absolute alcohol by heating in a boiling water-bath. After heating for 1 hour 30 minutes in the boiling water-bath, the reaction medium is poured into 300 g of iced water. The expected partially deacetylated product precipitates. It is filtered off, washed with water and dried in vacuo. It will be used as such for the second stage of the synthesis.

5

15

Second step

Preparation of 1-(2'-amino-5'-nitrophenoxy)-3-N-(β-hydroxyethyl)-aminopropan-2-ol 10 8.7 g of the crude product obtained in the first stage of the synthesis are heated for 30 minutes, 10

in a boiling water-bath, in 22 ml of hydrochloric acid (specific gravity=1.18). The cooled reaction mixture is treated with ammonia up to a pH of 6. The expected product precipitates in the form of the monohydrochloride. This monohydrochloride is filtered off, recrystallised from an ethanol/water 15 mixture and dried in vacuo. It melts at 156°C.

15

The monohydrochloride is dissolved in 20 ml of water. The aqueous solution is rendered alkaline with 20% strength ammonia solution. The 1-(2'-amino-5'-nitrophenoxy)-3-N-(β-hydroxyethyl)aminopropan-2-ol precipitates. The product is filtered off, washed with water, recrystallised from water and dried in vacuo. It melts at 177°C.

20

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₁ H ₁₇ N ₃ O ₅	Found
. C%	48.70	48.65
Н%	6.32	6.25
N%	15.49	15.44
0%	29.49	29.33

Preparation of 1-(2',5'-diaminophenoxy)-3-aminopropan-2-ol trihydrochloride

$$\begin{array}{c}
 & \text{NH}_2 \\
 & \text{O-CH}_2\text{-CHOH-CH}_2\text{-NH}_2 \\
 & \text{NH}_2
\end{array}$$

$$\begin{array}{c}
 & \text{O-CH}_2\text{-CHOH-CH}_2\text{NH}_2 \\
 & \text{3HC1}
\end{array}$$

7.5 g of zinc powder and a solution of 0.3 g of ammonium chloride in 1.5 ml of water are added 25 to 15 ml to 96° strength alcohol. The mixture is heated to the reflux temperature, with stirring, and 0.015 mol (3.4 g) of the 1-(2'-amino-5'-nitrophenoxy)-3-aminopropan-2-ol prepared according to Example 5 is then added gradually so as to maintain the reflux without external heating. When the addition has ended, the reflux is maintained for 5 minutes and the boiling reaction medium is then filtered into 4.6 ml of Iced hydrochloric acid (specific gravity=1.18). The expected product crystallises 30 in the form of the trihydrochloride. The product is filtered off, washed with alcohol and dried in vacuo. It melts with decomposition at 216-218°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for $C_9H_{15}N_3O_2$. HCI	Found	
C%	35.25	34.97	
Н%	5.92	5.86	
N%	13.70	13.82	
0%	10.43	10.68	
CI%	34.69	34.52	

5

10

Exampl 8
Preparation of 1-(2',4'-diamin phenoxy)-3-aminopropan-2-ol trihydrochl ride

0.02 mol (5.3 g) of the 1-(2'-amino-4'-nitrophenoxy)-3-aminopropan-2-ol monohydrochloride prepared according to Example 4 is introduced into 50 ml of 96° strength alcohol and 0.2 g of 10% strength palladium-on-charcoal. The mixture is heated at 85°C for one hour under 35 bars of hydrogen. After cooling, it is filtered to remove the catalyst, the filtrate being collected in 20 ml of iced ethanol saturated with hydrogen chloride. The expected product precipitates in the form of the trihydrochloride. After recrystallisation of this trihydrochloride from an ethanol/water mixture, it melts with decomposition at 236—238°C.

The product obtained corresponds to the following analysis results:

. Analysis	Calculated for C ₉ H ₁₅ N ₃ O ₂ . 3HCl	Found	
С%	35.23	35.41	
H%	5.87	5.90	
N%	13.70	13.63	
0%	10.44	10.63	
CI%	34.75	34.85	

It has been found, surprisingly, that this compound, which constitutes a coupler, gives, on coupling with para-aminophenol, much redder tints than its homologue which does not contain an OH group on the side chain; it has also been found, surprisingly, that the red tints obtained are much more stable than in the case of the non-hydroxylated homologue.

Exampl 9 Preparation of 1-(3'-methylamino-4'-nitrophenoxy)-3-aminopropan-2-ol hydrochloride

First step

5 5 Preparation of 1-(3'-methylamino-4'-nitrophenoxy)-2,3-epoxypropane 0.15 mol (36.3 g) of 3-N-methylamino-4-nitrophenyl β , γ -dihydroxypropyl ether (product described in Example 2 of French Patent Application 80/17,617) is dissolved in 150 ml of pyridine at ambient temperature. 0.22 mol (25.2 g) of methanesulphonyl chloride is added dropwise, in the course of 10 minutes and with stirring, to this pyridine solution kept at -5°C. When the addition has ended, 10 the temperature of the reaction medium is allowed to rise to about 20°C, and then, after the reaction 10 medium has been left to stand for 45 minutes at ambient temperature, 100 ml of 27.8% strength methanolic solution of sodium methylate are added thereto, at -5°C, in the course of 10 minutes. When the addition has ended, the mixture is stirred for 2 hours at ambient temperature and the methylene chloride is then washed with water, with 1 N hydrochloric acid solution and then with a 15 saturated solution of sodium bicarbonate. The methylene chloride is driven off in vacuo. The expected 15 product, consisting of the residual oil, crystallises very rapidly. After recrystallisation from ethyl acetate and drying in vacuo, it melts at 113°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₀ H ₁₂ N ₂ O ₄	Found	
. С%	53.57	53.66	
H%	5.39	5.35	
N%	12.50	12.48	
0%	28.54	28.58	

15

20

Second st p

Preparation f1-(3'-methylamino-4'-nitrophenoxy)-3-succinimid propan-2-ol

O.087 mol (19.5 g) of 1-(3'-m thylamino-4'-nitrophenoxy)-2,3-epoxypropane is dissolved in 75 ml of abs lut alcohol to which 6 dr ps of pyridine have been added. 0.10 mol (10.1 g) of succinimide is added and the mixture is then heated under reflux for 4 hours. The reaction mixture is filtered at the boil. On cooling of the filtrate, the expected product crystallises. After recrystallisation from alcohol and drying in vacuo, it melts at 152°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₄ H ₁₇ N ₃ O ₆	Found	
C%	52.01	51.97	
Н%	5.30	5.36	
N%	13.00	12.98	
0%	29.69	29.50	

10 Third step

Preparation of 1-(3'-methylamino-4'-nitrophenoxy)-3-aminopropan-2-ol monohydrochloride
10.5 g (0.0324 mol) of the succinimido derivative obtained according to the second step are
heated under reflux for 7 hours in 20 ml of 96° strength alcohol and 60 ml of 36% strength
hydrochloric acid. The reaction medium is cooled to 0°C. The expected product crystallises. It is filtered
off, washed with acetone and recrystallised from a mixture of water and alcohol. After drying in vacuo,
it melts with decomposition at between 258 and 260°C.

The product obtained corresponds to the following analysis results:

Analysis	Calculated for C ₁₀ H ₁₅ N ₃ O ₄ . HCl	Found
С%	43.25	43.21
Н%	5.81	5.78
N%	15.13	15.11
0%	23.04	23.10
CI%	12.77	12.70

Example 10
20 Preparation of 1-(2',4'-diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride mono-hydrate

10

. 15

20

25

15

First step

Preparati n of 1-(2'-amino-4'nitrophen xy)-3-dim thylaminopropan-2-ol

0.085 mol (21.5 g) of 1-(2'-ac tylamino-4'-nitrophenoxy)-2,3-epoxypropane (compound described in Example 4) is introduc d into 100 ml of a 40% strength aqueous solution of dimethylamine, and the reaction mixture is stirred for 30 minutes at ambient temperature. After dilution by the addition of 225 ml of iced water, followed by neutralisation with acetic acid, the mixture is evaporated to dryness in vacuo. The dried residue is taken up in 150 ml of ethyl acetate. After washing with acetone, the material which is insoluble in ethyl acetate consists of 28.5 g of 1-(2'-acetylamino-4'-nitrophenoxy)-3-dimethylaminopropan-2-ol. This acetylated derivative is heated under reflux for two hours in 220 ml of ethanol saturated with hydrogen chloride. After the reaction medium has cooled, the expected product in the form of the hydrochloride is filtered off. After this hydrochloride has been dissolved in water and the aqueous solution has been rendered alkaline with sodium hydroxide solution, the 1-(2'-amino-4'-nitrophenoxy)-3-dimethylaminopropan-2-ol is precipitated in the form of crystals.

The product is filtered off, washed with water and dried in vacuo. It melts at 123°C. Elementary analysis gives the following results:

Analysis	Calculated for C ₁₁ H ₁₇ N ₃ O ₄	Found
С%	51.76	51.65
Н%	6.67	6.72
N%	16.47	16.40
0%	25.10	25.02

Second step
Preparation of 1-(2',4'-diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride monohydrate

A solution of 0.12 mol (30.6 g) of 1-(2'-amino-4'-nitrophenoxy)-3-dimethylaminopropan-2-ol in 92 ml of absolute alcohol is subjected to catalytic hydrogenation at 80°C, under a pressure of 40 bars of hydrogen, in the presence of 4.5 g of 10% strength palladium-on-charcoal. The catalyst is removed by filtration, the alcoholic filtrate being received in 143 ml of iced ethanol saturated with hydrogen chloride. The expected product precipitates in the form of the trihydrochloride. This trihydrochloride is filtered off and recrystallised from an aqueous-alcoholic solution of hydrochloric acid. After drying in vacuo, it melts with decomposition at between 205 and 206°C.

Elementary analysis gives the following results:

Analysis	Calculated for $C_{11}H_{19}N_3O_2$. 3HCI. H_2O	Found
. C%	37.44	37.31
Н%	6.81	6.87
N%	11.91	11.79
. 0%	13.62	13.89
CI%	30.21	30.07

15

20

Example 11
Preparation of 2-hydroxy-3-(2',4'-diamin phenoxy)-propyltrimethylammonium chlorid dihydr chloride hemihydrate

5 First step
Preparation of 1-(2',4'-diacetylaminophenoxy)-3-dimethylaminopropan-2-ol

NH₂

Under the conditions described in the second step of Example 10, an alcoholic solution of 0.1 mol (25.5 g) of 1-(2'-amino-4'-nitrophenoxy)-3-dimethylaminopropan-2-ol is subjected to catalytic hydrogenation in the presence of palladium-on-charcoal. The catalyst is removed by filtration, the solvent is driven off in vacuo, the oily residue is taken up in 200 ml of ethyl acetate, and 17 ml of acetic anhydride are added gradually thereto. The expected diacetylated derivative precipitates in the form of crystals. It is filtered off, washed with acetone and dried in vacuo. It melts at 118—120°C.

Second step

Preparation of 2-hydroxy-3-(2',4'-diacetylaminophenoxy)-propyltrimethylammonium iodide monohydrate

0.03 mol (9.3 g) of the diacetylated derivative obtained in the first step is dissolved in 30 ml of acetone to which 3 ml of water has been added. The solution is heated to 50°C and 0.06 mol (3.75 ml) of methyl iodide is then added thereto, with stirring. The reaction medium is kept at 50°C for 30 minut s. The expected quaternary derivative precipitates. It is filtered off, washed with acetone and dried. It melts with decomposition at between 159 and 160°C.

Elementary analysis gives the following results:

Analysis	Calculated for C ₁₆ H ₂₆ O ₄ N ₃ J . H ₂ O	Found
C%	40.94	40.87
Н%	5.97	6.02
N%	8.95	8.87
0%	17.06	17.21
- 1%	27.08	27.02

Third step

Preparation of 2-hydroxy-3-(2',4'-diaminophenoxy)-propyltrimethylammonium dihydrochloride

5 hemihydrate

10

0.0137 mol (6.2 g) of the quaternary salt obtained in the second step is heated under reflux for 3 hours in 30 ml of ethanol saturated with hydrogen chloride. The expected product precipitates. It is filtered off, washed with absolute alcohol and dried in vacuo. It melts with decomposition at between 258 and 260°C.

Elementary analysis gives the following results:

5

10

30

Analysis	Calculated for $C_{12}H_{22}N_3O_2CI$. 2HCI. 1/2H ₂ O	Found
С%	40.28	40.51
H%	6.99	6.72
N%	11.75	11.77
0%	11.19	11.21
CI	29.80	29.95

Example 12

The following dyeing composition is prepared:

	1-(3'-Nitro-4'-aminophenoxy)-3-aminopropan-2-ol	. 0.2 g	45
15	2-Butoxyethanol	10 g	15
	Hydroxyethylcellulose	2 g	
	Ammonium lauryl-sulphate	5 g	
	Water qs	100 g	
The pH of the composition is equal to 8.4. When applied to bleached hair for 20 minutes at 30°C, this mixture imparts to the hair, after rinsing and shampooing, a colouration of 7 YR 7/14 (determined on the "Munsell" scale).		re imparts to the hair, after "Munsell" scale).	20
	Example 13 The following dyeing composition is prepared:		

1-(2'-Amino-4'-nitrophenoxy)-3-aminopropan-2-ol 25 2-Butoxyethanol	. 0.8 g 10 g	25
Acrylic acid polymer having a molecular weight of 2 to 3 million, solder under the name "Carbopol 934" by "Goodrich Chemical Co." Ammonia solution (22°Bé strength) Water qs	2 g 2 g 100 g	

The pH of the composition is equal to 6.7.

When applied to bleached hair f r 35 minutes at 30°C, this mixture imparts to the hair, after rinsing and shampooing, a colourati n of 3.75 Y 8.5/9.5 (determined on the "Munsell" scale).

	Example 14		
	The following dy ing composition is prepared:	•	
	1-(3'-Nitro-4'-aminophen xy)-3-N,N-di thylaminopropan-2-ol		
	hydrochloride	0.6 g	
5	Propylene glycol	10 g	5
	Diethanolamides of copra fatty acids	2.2 g	
	Lauric acid	0.8 g	
	Ethylene glycol monoethyl ether	2 g	
	Monoethanolamine	1 g	_
10	Water qs	100 g	10
	The pH of the composition is equal to 7.8.		
	When applied to bleached hair for 25 minutes at 30°C, this mixture impa		
	rinsing and shampooing, a colouration of 7.5 YR 7/16 (determined on the "Mu	ilseli scale).	
	Example 15		15
15	The following dyeing composition is prepared:		
	1-(2'-Amino-5'-nitrophenoxy)-3-aminopropan-2-ol	0.55 g	
		10 g	•
	Propylene glycol Lauric acid monoethanolamide	1.5 g	
	Lauric acid	1 g	
	Hydroxyethylcellulose	5 g	20
20	Monoethanolamine	2 g	
		100 g	
	Water qs		
	The pH of the composition is equal to 10.		•
	When applied to bleached hair for 25 minutes at 25°C, this mixture impa	arts to the hair, after	
25	rinsing and shampooing, a colouration of 6.25 Y 8.5/12 (determined on the "N	/lunsell" scale).	25
25	Thising the shampooning, a coloured control of the colour	· ·	•
	Example 16		
	The following dyeing composition is prepared:		
	The teneving Lyong temperature (
	1-(3'-Nitro-4'-aminophenoxy)-3-N-propylaminopropan-2-ol		
	monohydrochloride	1 g	
30	2-Butoxyethanol	10 g	30
-	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea"	8 g	
	Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by		
	"Henkel"	0.5 g	
	Oxyethyleneated castor oil sold under the name "Cemulsol B" by "Rhone-		
35	Poulenc"	1 <u>g</u>	35
-	Oleic diethanolamide	1.5 g	
	Ammonia solution (22°C Bé strength)	0.25 g	
	Water qs	100 g	
	The pH of the composition is equal to 8.7.		4.0
40	When applied to bleached hair for 35 minutes at 28°C, this mixture imp	arts to the hair, after	40
	rinsing and shampooing, a colouration of 7 YR 7/16 (determined on the "Mun	sell" scale).	
		•	
	Example 17		
	The following dyeing composition is prepared:		
		0.4 ~	
	1-(2'-Amino-5'-nitrophenoxy)-3-N-(β-hydroxyethyl)-aminopropan-2-ol	0.4 g	45
45	2-Butoxyethanol	10 g	40
	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea"	8 g	
	Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by	0 E -	
	"Henkel"	0.5 g	
	Oxyethyleneated castor oil s ld under the name "Cemulsol B" by "Rh ne-	4 -	E0
50		1 g	50
	Ol ic diethan lamide	1.5 g	•
	Triethanolamin (containing 10% factive ingredient)	0.4 g	
	Water qs ·	100 g	

The pH of the c mp sition is equal to 8.7.

23 GB 2 129 022 A

When applied to bleach d hair for 30 minutes at 28°C, this mixture imparts t the hair, after rinsing and shampooing, a colouration of 7.5 Y 8.5/11 (determined on the "Munsell" scale).

Exampl 18

	The following dy ing comp sition is prepared:		
5	1-(3'-Nitro-4'-aminophenoxy)-3-N-propylaminopropan-2-ol hydrochloride 3-Nitro-4-N'-methylamino-N-(β -aminoethyl)-aniline dihydrochloride 2-Butoxyethanol	0.15 g 0.076 g 10 g	.
	Hydroxyethylcellulose sold under the name "Cellosize WPO3" by "Union	_	
	- Carbide"	2 g	10
10	Dimethyl-alkyl-hydroxyethyl-ammonium bromide (alkyl=tallow derivative)	2 g 0.25 g	10
	Ammonia solution (22° Bé strength)	100 g	
	Water qs	100 8	
	The pH of the composition is equal to 8.5.	nivtura imparts to the	•
	When applied to 90% naturally white hair for 25 minutes at 28°C, this r	IIIXtule Impai & to aic	15
15	hair, after rinsing and shampooing, a coppery light chestnut colouration.		
	Example 19		
	The following dyeing composition is prepared:		
	1-(3'-Nitro-4'-aminophenoxy)-3-N,N-diethylaminopropan-2-ol	0.05 g	
	hydrochloride	0.05 g	20
20	3-Nitro-4-N'-(β-aminoethyl)-amino-N,N-di-(β-hydroxyethyl)-aniline	0.07 g	
	dihydrochloride 3-Nitro-4-amino-6-methyl-N-(β-hydroxyethyl)-aniline	0.03 g	
	2-Butoxyethanol	10 g	
	Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under		
25	the name "Cemulsol NP." by "Rhone-Poulenc"	12 g	25
	Nonyiphenol oxyethyleneated with 9 mols of ethylene oxide, sold under		
	the name "Cemulsol NP _o " by "Rhone-Poulenc"	15 g	
	Olevi alcohol oxyethyleneated with two mols of ethylene oxide	1.5 g	
	Oleyi alcohol oxyethyleneated with four mols of ethylene oxide	1.5 g	30
30	Triethanolamine (containing 20% of active ingredient)	1.5 g 100 g	30
	Water qs	100 8	
•	The pH of the composition is equal to 8.4.		
	When applied to 90% naturally white hair for 20 minutes at 28°C, this	mixture imparts to the	
	hair, after rinsing and shampooing, a pinkish beige colouration.		
			35
· 35	Example 20		
	The following dyeing composition is prepared:		•
	1-(2'-Amino-4'-nitrophenoxy)-3-aminopropan-2-ol	· 0.2 g	
	3-Nitro-4-N-(β -hydroxyethyl)-aminophenyl β -hydroxypropyl ether	0.08 g	
	3-Nitro-4-amino-N-(B-hydroxypropyl)-aniline	0.06 g	40
40	2-[4'-(N,N-dihydroxyethylamino)-anilino]-5-N'-(β-hydroxyethyl)-amino-		40
	1,4-benzoquinone	0.5 g	
	2-Butoxyethanol	10 g	
	Diethanolamides of copra fatty acids	2.2 g 0.8 g	
	Lauric acid	0.0 g 2 g	45
45	Ethylene glycol monoethyl ether Monoethanolamine	1 g	
	Water qs	100 g	
	•	<u> </u>	
	The pH of the composition is equal to 7.		
	When applied to bleached hair for 20 minutes at 30°C, this mixture im	parts to the hair, after	50
50	the state of the second	•	Ų.

The following dyeing composition is prepared:

	1-(3'-Nitro-4'-aminophenoxy)-3-N-propylaminopropan-2-ol		
		0.065 g	
	m nohydrochloride	0.035 g	55.
55	2-N-(β-Hydroxyethyl)-amino-5-nitrophen	0.035 g	99.

24		GB 2 129 022 A	24
		0.04	
	3-Nitro-4-N'-methylamino-N,N-di-(β-hydroxyethyl)-aniline	0.21 g	
	3-Nitro-4-amino-6-m thyl-N-(β,γ-dihydroxypr pyl)-aniline	0.05 g	
	2-But xy thanol	10 g 2 g	
E	Carboxymethylcellulos Ammonium lauryl-sulphate	5 g	5
9	Triethanolamine (containing 20% of active ingredient)	0.125 g	Ū
	Water qs	100 g	
	The pH of the composition is equal to 7.		
	When applied to bleached hair for 25 minutes at 30°C, this mixture impa	arts to the hair, after	
10	rinsing and shampooing, a golden sandy colouration.		10
	Example 22		
	The following dyeing composition is prepared:		
	A TOUR BY ALL IN TO A STATE OF THE STATE OF	0.055 -	
	1-(3'-Nitro-4'-aminophenoxy)-3-aminopropan-2-ol	0.055 g	٠.
	1,4,5,8-Tetraaminoanthraquinone 2-Amino-3-nitrotoluene	0.105 g 0.035 g	15
15	2-Amino-3-introtoluene 3-Nitro-4-amino-6-methyl-N-(β-aminoethyl)-aniline	0.033 g 0.03 g	13
	Propylene glycol	10 g	
	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea"	8 g	
	Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by	J	
20	"Henkel"	0.5 g	20
	Oxyethyleneated castor oil sold under the name "Cemulsol B" by "Rhone-		•
	Poulenc"	1 g	
	Oleic diethanolamide	1.5 g	
	Triethanolamine (containing 20% of active ingredient)	2 g	٥٢
25	Water qs	100 g	25
,	The pH of the composition is equal to 9.	,	
	When applied to 90% naturally white hair for 40 minutes at 28°C, this n	nixture imparts to the	
	hair, after rinsing and shampooing, a golden blond colouration.		
	- 1.00		
	Example 23		30
30	The following dyeing composition is prepared:		
	1-(2'-Amino-5'-nitrophenoxy)-3-aminopropan-2-ol	0.1 g	
	3-Nitro-4-N-(β-aminoethyl)-aminophenoxyethanol	0.15 g	
	3-Nitro-4-amino-6-methyl-N-(β,γ-dihydroxypropyl)-aniline	0.05 g	
	3-Nitro-4-N'-methylamino-N,N-di-(β-hydroxyethyl)-aniline	0.5 g	
35	Propylene glycol	10 g	35
	Hydroxyethylcellulose sold under the name "Cellosize WPO3" by "Union	_	
	Carbide"	2 g	
	Dimethyl-alkyl-hydroxyethyl-ammonium bromide (alkyl=tallow derivative)	2 g	
	Ammonia solution qs	pH 8	40
40	Water qs · · · · · · · · · · · · · · · · · ·	100 g	
	When applied for 25 minutes at 28°C to hair which has been bleached		•
	imparts to the hair, after rinsing and shampooing, a very coppery chestnut col	ouration.	
	Enamela 24		
	Example 24 The following dyeing composition is prepared:		
	the following dyeing composition is prepared:	•	
45	1-(3'-Nitro-4'-aminophenoxy)-3-aminopropan-2-ol	0.5 g	45
40	1-(2'-Amino-5'-nitrophenoxy)-3-aminopropan-2-ol	0.2 g	
	3-Nitro-4-amino-N-(β-hydroxyethyl)-aniline	0.15 g	
	3-Nitro-2-N'-(β-aminoethyl)-amino-N,N-di-(β-hydroxyethyl)-aniline	• .	
	dihydrochloride	0.2 g	
50	Hydroxyethylcellulose	2 g	50
•	Ammonium lauryl-sulphat	5 g	
	Ammonia solution (22° Bé strength)	0.25 g	
•	Water qs ·	100 g	•
	The all afab assessed to the second of O		
-	The pH of the composition is equal to 9. When applied to bleached hair for 20 minutes at 28°C, this mixture imp	arte to the hoir often	66
55	rinsing and shampo ing, a reddish copper c louration.	ज कि (118 Hall, व्राप्ति	55
	mong and anampo mg, a readion opper o toutation	•	

	·	•	
	Example 25		
	The following dyeing composition is prepared:		
	1-(2',4'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.11 g	
	Resorcin 1	0.08 g	
5	Para-phenylenediamine	0.3 g	5
•	4-Amino-N,N-di-(β-hydroxyethyl)-aniline dihydrochloride	0.6 g	
	3-Nitro-4-amino-N-(β-hydroxypropyl)-aniline	0.5 g	
	3-N-Methylamino-4-nitrophenoxyethanol	0.3 g 8 g	
40	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea" Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by	O g	10
10	"Henkel"	. 0.5 g	
	Oxyethyleneated castor oil sold under the name "Cemulsol B" by "Rhone-		•
	Poulenc"	1 g	
	Oleic diethanolamide	1.5 g	15
15	Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the	2.5 g	13
•	name "Masquol DTPA" Sodium bisulphite solution (35° Bé strength)	1 g	
•	Ammonia solution (22° Bé strength)	11 g	
	Water qs	100 g	
			20
20	The pH of the composition is equal to 10.1.	me of use	20
	100 g of hydrogen peroxide of 20 volumes strength are added at the ti When applied for 30 minutes at 30°C to hair which has been bleached	I straw vellow, this mixture	
	imparts to the hair, after rinsing and shampooing, a black-brown colouration		
	impures to the hair, arter thining and enempeering,	•	
	Example 26		
25	The following dyeing composition is prepared:	·	25
	1-(2',4'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.4 g	
	Resorcinol	0.1 g	
	Meta-aminophenol	0.13 g	
	Para-phenylenediamine	0.125 g	20
30	Para-aminophenol	0.13 g	30
	N-Methyl-para-aminophenol sulphate	0.3 g 0.4 g	
	2-Methyl-4-amino-5-nitrophenol Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under	0.4 g	
	the name "Remcopal 334" by "Gerland"	. 21 g	
35	Nonylphenol oxyethyleneated with 9 mols of ethylene oxide, sold under		35
	the name "Remcopal 349" by "Gerland"	24 g	
	Oleic acid	4 g 3 g	
	2-Butoxyethanol	10 g	
40	Ethanol (96° strength) Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the	. o g	40
40	· name "Masquol DTPA"	2.5 g	
	Sodium bisulphite solution (35° Bé strength)	1 g	
	Ammonia solution (22° Bé strength)	10 g	
	Water qs	100 g	
AC	The pH of the composition is equal to 10.5.	·	45
45	75 g of hydrogen peroxide of 20 volumes strength are added at the tir	ne of use.	
	When applied to 90% naturally white hair for 25 minutes at 30°C, this	s mixture imparts to the	
	hair, after rinsing and shampooing, an ashen chestnut colouration.		
	Example 27 The following dyeing composition is prepared:		50
50	The following dyeing composition is prepared.		
	1-(2',5'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.153 g	
	2 6-Dimethyl-3-acetylaminophenol	0.089 g	
	Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under	21.5	
	the name "Remcopal 334" by "G rland"	. 21 g	55
55	Nonylphenol oxyethyleneated with 9 mols of ethylene oxide, sold under the name "R mcopal 349" by "Gerland"	24 g	55
	Oleic acid	4 g	
	2-Butoxyethanol	3 g	
			•

GB 2 129 022 A 26

			
	THE MODE 1 111	10 g	
	Ethanol (96° strength) Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the	. IOG	
		3.5 g	
	nam "Masquol DTPA" Ammonia solution (22° Bé strength)	.10 g	
5	Water as	100 g	5
. 3	vvacei q5	1009	٠. •
	The pH of the composition is equal to 10.5. 100 g of hydrogen peroxide of 20 volumes strength are added at the tim When applied to 90% naturally white hair for 25 minutes at 30°C, this n hair, after rinsing and shampooing, a bluish grey colouration.	e of use. nixture imparts to the	
			40
10	Example 28 The following dyeing composition is prepared:		10
	1-(2',5'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.63 g	
	2-Methylresorcinol	0.14 g	
	3-Acetylaminophenol	0.135 g	
15	2,4-Diaminophenoxyethanol dihydrochloride	0.06 g	15
13	3-Nitro-4-amino-6-methyl-N-(β-hydroxyethyl)-aniline	0.215 g	
	1-(3'-Nitro-4'-aminophenoxy)-3-aminopropan-2-ol	0.3 g	
	Acrylic acid polymer having a molecular weight of 2 to 3 million, sold		
	under the name "Carbapol 934" by "Goodrich Chemical Co."	1.5 g	
20	Alcohol (96° strength)	11 g	20
20	2-Butoxyethanol	5 g -	
	Trimethylcetylammonium bromide	1 g	
	Ethylenediaminetetraacetic acid sold under the name "Trilon B"	0.1 g	
	Ammonia solution (22° Bé strength)	10 g	
25	Thioglycolic acid	0.2 g	25
-	Water qs	100 g	
30	The pH of the composition is equal to 10. 100 g of hydrogen peroxide of 20 volumes strength are added at the tin When applied for 25 minutes at 30°C to hair which has been bleached imparts to the hair, after rinsing and shampooing, a chestnut colouration.	ne of use. straw yellow, this mixtu	ıre 30
	Example 29 The following dyeing composition is prepared:		
	The following dyoing composition to properties		
	1-(2',5'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.306 g	
	2.4-Diaminophenyl β-hydroxypropyl ether dihydrochloride	0.255 g	
35	Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under		35
••	the name "Cemulsol NP," by "Rhone-Poulenc"	12 g	
	Nonylphenol oxyethyleneated with 9 mols of ethylene oxide, sold under		
	the name "Cemulsol NP _a " by "Rhone-Poulenc"	15 g	
	Oleyl alcohol oxyethyleneated with two mols of ethylene oxide	1.5 ⁻ g	400
40	Oleyl alcohol oxyethyleneated with four mols of ethylene oxide	1.5 g	40
	Propylene glycol	6 g	
	Ethylenediaminetetraacetic acid sold under the name "Trilon B"	0.12 g	
	Ammonia solution (22° Bé strength)	11 g	
	Thioglycolic acid	0.6 g 100 g	45
45	Water qs	100 8	40
	The pH of the composition is equal to 9.9. 100 g of hydrogen peroxide of 20 volumes strength are added at the tir When applied to 90% naturally white hair for 30 minutes at 28°C, this hair, after rinsing and shampooing, a pure blue colouration.	ne of use. mixture imparts to the	
	Francis 20		50
50	Example 30 The following dyeing composition is prepared:		
	1-(2',4'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.25 g	•
	Para-ph nylenediamin	0.088 g	
	Oleyl alcoh I oxyethyleneated with two mols of ethylene oxide	4.5 g	
55	Oleyl alcohol oxyethyleneated with four mols of ethylen xide	4.5 g	55
-	Oleylamine oxyethyleneat d with 12 m ls f thylen oxid, sold und r		
	the nam "Ethom en TO ₁₂ " by "Armour Hess"	4.5 g	
	14		

27		GB 2 129 022 A	
	Diethanolamides of copra fatty acids	9 g	
	Propylene glycol	4 g	
	2-Butoxyethanol	8 g	
	Ethanol (96° strength)	6 g	
5	Pentasodium salt of diethylenetriaminepentaacetic acid, sold under th		5
J	name "Masquol DTPA"	2 g	
	Thioglycolic acid	0.5 g	
	Ammonia solution (22° Bé strength)	10 g	
		100 g	
	Water qs	.009	
40	The pH of the composition is equal to 10.3.		10
10	85 g of hydrogen peroxide of 20 volumes strength are added at the tim	ne of use.	10
	When applied to 90% naturally white hair for 25 minutes at 30°C, this	mixture imparts to the	
	tynen applied to 50% flaturally write flat for 25 fillings at 50°C, this	Minitare imparte to the	
	hair, after rinsing and shapooing, a midnight blue colouration.		
	Example 31		15
15	The following dyeing composition is prepared:		10
		0.12 -	
	1-(2',4'-Diaminophenoxy)-3-aminopropan-2-ol trihydrochloride	0.12 g	
	Para-aminophenol	0.043 g	_
	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea"	19 g	
	2-Octyldodecanol sold under the name "Eutanol G" by "Henkel"	4.5 g	
20	Cetyl/stearyl alcohol containing 15 mols of ethylene oxide, sold under the		20
	name "Mergital CS 15/E" by "Henkel"	2.5 g	
	Ammonium lauryl-sulphate containing 30% of active ingredient	12 g	
	Cationic polymer possessing repeat units:		
	$+ (CH_2)_2 - N (CH_2)_5 +$	4 g	
	$ \left\{ \begin{array}{c} \bigoplus_{\text{C1} \ominus \text{CH}_3} (\text{CH}_2)_3 - \bigvee_{\text{C1} \ominus \text{CH}_3} (\text{CH}_2)_6 \\ \end{array} \right\} $		
		2 g	25
25	Benzyl alcohol	11 g	
	Ammonia solution (22° Bé strength)	1 g	
	Ethylenediaminetetraacetic acid solid under the name "Trilon B"	1.2 g	
	Sodium sulphite	100 g	
	Water qs ·	100 g	
	m	• •	30
30	The pH of the composition is equal to 10.	na of use	•
٠	75 g of hydrogen peroxide of 20 volumes strength are added at the tir	nnarte to the hair after	
	When applied to bleached hair for 20 minutes at 28°C, this mixture in	inparts to the nam, area	
	rinsing and shampooing, a pink champagne colouration.		
	Everanie 22		
25	Example 32 The following dyeing composition is prepared:		35
35	The following dyeing composition to property	•	
	1-{2',4'-Diaminophenoxy}-3-aminopropan-2-ol trihydrochloride	0.05 g	
		0.15 g	
	Resorcinol	0.085 g	
	Meta-aminophenol	0.09 g	
	2-Methyl-5-N-(β-hydroxyethyl)-aminophenol	0.1 g	40
40	Para-phenylenediamine	0.405 g	
	2-Amino-3-nitrophenol	0.08 g	
	3-Nitro-4-amino-6-methyl-N-(β-aminoethyl)-aniline	0.00 g	
	Acrylic acid polymer having a molecular weight of 2 to 3 million, sold	1.5 g	
	under the name "Carbopol 934" by "Goodrich Chemical Co."	•	45
45	Alcohol (96° strength)	11 g	45
	2-Butoxyethanol	5 g	
	Trimethylcetylammonium bromide	1 g	
	Ethyl nediaminetetraacetic acid sold under the name "Trilon B"	0.1 g	
	Ammonia soluti n (22° Bé strength)	10 g	
50	Thioglycolic acid	0.2 g	50
30	Water qs	100 g	
	ereces of		

The pH of the composition is equal to 10.3. 100 g $\,$ f hydrog in peroxid of 20 volumes strength are add d at the time of us $\,$.

3 g

10 g

2.5 g

0.6 g

10 g

100 g

45

50

When appli d for 30 minutes at 28°C to hair which has be n bleach d straw yell w, this mixture imparts to the hair, after rinsing and shampooing, a r ddish copper m dium chestnut colouration.

Example 33

The following dyeing composition is prepared:

_	4 /2/ Mathulamina // nitranhanaus/ 2 aminanyanan 2 al		5
5	1-(3'-Methylamino-4'-nitrophenoxy)-3-aminopropan-2-ol	0.3 g	5
	monohydrochloride	0.3 g 0.8 g	
	3-Nitro-4-amino-6-methyl-N-(β-hydroxyethyl)-aniline		
	1,4,5,8-Tetraaminoanthraquinone	0.06 g	
	2-Butoxyethanol	10 g	
1.0	Hydroxyethylcellulose sold under the name "Cellosize WPO3" by "Union		10
	Carbide"	2 g	
	Dimethyl-alkyl-hydroxyethyl-ammonium bromide (alkyl=tallow derivative)	2 g	
	Ammonia solution (5% strength)	1 g	
	Water qs	100 g	
15	The pH of the composition is equal to 7.		15
	When applied to bleached hair for 20 minutes at 30°C, this mixture impa	irts to the hair, after	
	rinsing and shampooing, a light copper colouration.		
	E		
	Example 34		
	The following dyeing composition is prepared:		
20	1-(3'-Methylamino-4'-nitrophenoxy)-2-aminopropanol		20
20	monohydrochloride	1 g	
	2-Butoxyethanol	10 g	
	Cetyl/stearyl alcohol sold under the name "Alfol C16/18" by "Condea"	8 g	
	Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by	O 9	
25	"Henkel"	0.5 g	25
25	Oxyethyleneated castor oil sold under the name "Cemulsol B" by "Rhone-	0.5 g	23
	Poulenc"	1 a	
		1 g 1.5 g	
	Oleic diethanolamide		
	Triethanolamine (in 20% strength aqueous solution)	2 g	20
30	Water qs	100 g	30
	The AU of the composition is accord to 7.5		
	The pH of the composition is equal to 7.5. When applied to bleached hair for 30 minutes at 28°C, this mixture imparts.	arte to the bair after	
	rinsing and shampooing, a colouration of 4.5 Y 8.5/13 (determined on the "M	uncell" ecole)	
	mising and stampooling, a colouration of 4.5 1 6.5/15 (determined on the 14)	unach adalej.	
	Example 35		35
35	The following dyeing composition is prepared:		35
	1-(2',4'-Diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride	0.000	
	monohydrate	0.836 g	
	Para-phenylenediamine	0.256 g	
	Nonylphenol containing 4 mols of ethylene oxide, sold by "Rhone-		
40	Poulenc" under the name "Cemulsol NP4"	21 g	40
	Nonylphenol containing 9 mols of ethylene oxide, sold by "Rhone-		
	Poulenc" under the name "Cemulsol NP9"	24 g	
	Oleic acid	4 g	
		2	

Th pH of the composition is equal to 10.2.

Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the

2-Butoxyethanol 45 Ethanol (96° strength)

Thioglycolic acid

50 Water qs

name "Masquol DTPA"

Ammonia solution (22° Bé strength)

120 g f hydrogen peroxide of 20 volumes strength are add d at the time of us .

When applied to 90% naturally whit hair for 25 minutes at 28°C, this mixture imparts to the hair, after rinsing and shampooing, a midnight blue colouration.

	Example 36		
	The following dyeing composition is prepared:		٠.
	2-Hydroxy-3-(2',4'-diaminophenoxy)-propyltrimethylammonium chloride	0.05	
	dihydrochl ride h mihydrate	0.35 g 0.108 g	5
5	Para-phenylenediamine Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under	0.100 g	J
	the name "Remcopal 334" by "Gerland"	21 g	
	Nonylphenol oxyethyleneated with 9 mols of ethylene oxide, sold under	. •	•
	the name "Remcopal 349" by "Gerland"	. 24 g	10
10	Oleic acid	4 g 3 g	10
	2-Butoxyethanol	. 10 g	
	Ethanol (96° strength) Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the	, , , , g	
	· name "Masquol DTPA"	2.5 g	45
15	Sodium bisulphite solution (35° Bé strength)	1 g	15
	Ammonia solution (22° Bé strength)	10 g 100 g	
	Water qs	100 g	
	The pH of the composition is equal to 10.	,	
	100 g of hydrogen peroxide of 20 volumes strength are added at the time	e of use.	20
20	When applied to 90% naturally white hair for 20 minutes at 30°C, this r	nixture imparts to the	20
	hair, after rinsing and shampooing, a bluish grey colouration.		
	Example 37 The following dyeing composition is prepared:		
	The following dyeing composition is prepared:		
	1-(2',4'-Diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride		
25	monohydrate	0.35 g	25
	Para-aminophenol	0.109 g	
	Acrylic acid polymer having a molecular weight of 2 to 3 million, sold under the name "Carbopol 934" by "Goodrich Chemical Co."	3 g	
	Ethanol (96° strength)	11 g	
30	2-Butoxyethanol	5 g	30
	Trimethylcetylammonium bromide	2 g 0.2 g	
	Ethylenediaminetetraacetic acid sold under the name "Trilon B" Ammonia solution (22° Bé strength)	10 g	
	Sodium bisulphite solution (35° Bé strength)	1 g ັ	
35	Water qs	100 g	35
-			
•	The pH of the composition is equal to 9.8. 100 g of hydrogen peroxide of 20 volumes strength are added at the tir	ne of use.	
	When applied for 25 minutes at 28°C to hair which has been bleached	white, this mixture	
	imparts to the hair, after rinsing and shampooing, a pink shade with a golden	sheen.	
40	Example 38		40
	The following dyeing composition is prepared:		
	1-(2',4'-Diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride		
	monohydrate	0.13 g	
	Paraphenylenediamine	0.08 g	
45	Para-aminophenol	0.16 g 0.13 g	45
	Resorcinol	0.13 g 0.09 g	
	Meta-aminophenol 2-Methyl-5-N-(β-hydroxyethyl)-aminophenol	0.1 g	
	3-N-Methylamino-4-nitrophenoxyethanol	0.03 g	
50	Oleyl alcohol oxyethyleneated with 2 mols of ethylene oxide	4.5 g	50
	Olevi alcohol oxyethyl, neated with 4 mols of ethylene oxid	4.5 g	
	Oleylamin oxyethylen ated with 12 m is of ethylene oxide, sold under	4.5 g	
	the name "Ethomeen TO ₁₂ " by "Armour Hess" Diethanolamides of copra fatty acids	9 g	
EE	Propylene glycol	4 g	55
55	2-Butoxyethanol	8 g	
	Ethanol (96° strength)	6 g	

	Pentasodium salt of diethylenetriamin p ntaacetic acid, sold under the		
	name "Masqu I DTPA"	2 g	
	Hydroquinone	0.15 g	
	Sodium bisulphite solution (35° Bé strength)	1.3 g	
5	Ammonia solution (22° Bé strength)	10 g	5
	Water qs '-	100 g	_
	770.07 40		
	The pH of the composition is equal to 10.		
	120 g of hydrogen peroxide of 20 volumes strength are added at the time	e of use.	
·	When applied to 90% naturally white hair for 25 minutes at 28°C, this m		
10	hair, after rinsing and shampooing, a hazel colouration.	•	· 10
	Example 39		
	The following dyeing composition is prepared:	•	
	1-(2',4'-Diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride		_
	monohydrate	0.25 g	
	Para-phenylenediamine	0.4 g	15
	Para-aminophenol .	0.6 g	
	2-Methylresorcinol	0.4 g	
	2-Methyl-5-N-(β-hydroxyethyl)-aminophenol	0.25 g	
	2-Amino-3-nitrophenol	0.2 g	·
20	Nonylphenol oxyethyleneated with 4 mols of ethylene oxide, sold under		20
	name "Remcopal 334" by "Gerland"	21 g	
	Nonylphenol oxyethyleneated with 9 mols of ethylene oxide, sold under	0.4	
	the name "Remcopal 349" by "Gerland"	24 g	
	Oleic acid	4 g	
25	2-Butoxyethanol	3 g	25
	Ethanol (96° strength)	10 g	
	Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the	3 F -	
	name "Masquol DTPA" :	2.5 g	
	Sodium bisulphite solution (35° Bé strength)	1 g 10 g	
30	Ammonia solution (22° Bé strength)	100 g	30
	Water qs	100 g	
	The pH of the composition is equal to 10.1.		
	120 g of hydrogen peroxide of 20 volumes strength are added at the tim	e of use.	
	When applied for 15 minutes at 30°C to hair which has been bleached s		
35	imparts to the hair, after rinsing and shampooing, a coppery chestnut colourat		35
55	impurate to the many area, missing and arrange and arrange area.		-
	Example 40		
	The following dyeing composition is prepared:		
	1-(2',4'-Diaminophenoxy)-3-dimethylaminopropan-2-ol trihydrochloride		•
	monohydrate	0.075 g	
40	Para-phenylenediamine	0.20 g	40
70	N,N-Di-(β-hydroxyethyl)-paraphenylenediamine dihydrochloride	0.20 g	
	N-Methyl-para-aminophenol sulphate	0.115 g	
	2-Methylresorcinol	0.3 g	
	Meta-aminophenol	0.085 g	
45	1-Amino-2-nitro-4-N-(β-hydroxyethyl)-amino-5-methylbenzene	0.18 g	45
	3-Nitro-4-N-(β-aminoethyl)-aminophenoxyethanol	0.15 g	
	Cetyl/stearyl alcohol sold under the name "Alfol C16/18E" by "Condea"	8 g	
	Sodium cetyl/stearyl-sulphate sold under the name "Lanette Wax E" by	-	
	"Henkel"	0.5 g	
50			50
	Poulenc"	1 g	
	Oleic diethanolamide	1.5 g	
	Pentasodium salt of diethylenetriaminepentaacetic acid, sold under the		
	name "Masquol DTPA"	2.5 g	
55		0.3 g	55
	Ammonia s lution (22° Bé strength)	11 g	
	Wat rqs	100 g	

The pH of the c mposition is equal to 9.3. 100 g of hydrogen p roxide of 20 volumes strength are added at th tim of use.

When applied to 90% naturally whit hair for 25 minutes at 25°C, this mixtur imparts to the hair, after rinsing and shampooing, a d ep chestnut colouration with a c ppery sheen.

Example 41

Th following dyeing compositi n is prepared:

5	2-Hydroxy-3-(2',4'-diaminophenoxy)propyltrimethylammonium chloride		5
	dihydrochloride hemihydrate	0.1 g	
	Para-toluylenediamine dihydrochloride	0.085 g	
	Para-aminophenol	0.155 g	
	Resorcinol	0.08 g	
10	Meta-aminophenol	0.09 g	10
	2-Methyl-5-N-(β-hydroxyethyl)-aminophenol	0.1 g	
	Acrylic acid polymer having a molecular weight of 2 to 3 million, sold		
	under the name "Carbopol 934" by "Goodrich Chemical Co."	1.5 g	
	Ethanol (96° strength)	11 g	
15	2-Butoxyethanol :	5 g	15
	Trimethylcetylammonium bromide	1 g	
	Ethylenediaminetetraacetic acid sold under the name "Trilon B"	0.1 g	
	Ammonia solution (22° Bé strength)	10 g	
	Thioglycolic acid	0.2 g	
20	Water qs	100 g	20

The pH of the composition is equal to 9.9.

100 g of hydrogen peroxide of 20 volumes strength are added at the time of use.

When applied to 90% naturally white hair for 25 minutes at 28°C, this mixture imparts to the hair, after rinsing and shampooing, a hazel colouration.

It is clearly understood that the embodiments described above in no way imply a limitation and can form the subject of any desirable modifications without thereby exceeding the scope of the invention.

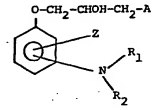
Claims

25

1. A dyeing composition suitable for keratin fibres and in particular for hair, which composition comprises at least one compound, or salt thereof, of the formula (I)

30

25



in which formula:

Z represents a nitro group; A represents a group NY or

> ø/ N

35

wherein

35

Y represents two identical or different C_1 — C_4 alkyl or C_1 — C_4 hydroxyalkyl substituent groups or together with the nitrogen atom of the group NY or



represents a morpholino or piperidino heterocyclic ring or, when A represents the group NY, may also represent two hydrogen atoms or a hydrogen atom and a C₁—C₄ alkyl or C₁—C₄ hydroxyalkyl substituent group, Y' represents a C₁—C₄ alkyl substituent group and X r presents an anion; and

41.

	R_1 and R_2 are identical or different and each represents a hydrogen atom or a C_1 — C_4 alkyl or C_1 — C_4 hydroxyalkyl radical; together with an appropriate carrier.	
5	2. A dyeing composition suitable for k ratin fibers and in particular for hair, which composition comprises at least one compound of the formula (I), or salt thereof, as defined in claim 1 except that Z represents an NH ₂ group in the ortho- or para-position relative to NR ₁ R ₂ , together with an appropriate	5
	3. A composition according to claim 2 which contains an oxidising agent at the time of use. 4. A composition according to any one of claims 1 to 3, wherein the compound of the formula (I)	10
10	is present in the composition in an amount of from 0.001 to 4% by weight, relative to the total weight of the composition.	10
	5. A dyeing composition suitable for keratin fibres and in particular for hair, which composition comprises at least one compound of the formula (I), or salt thereof, as defined in claim 1 except that Z represents an NH ₂ group in the meta-position relative to NR ₁ R ₂ , and if A denotes NY, if Z is in the 4-	
15	position on the benzene nucleus and if NR_1R_2 is in the 2-position, R_1 and R_2 both representing a hydrogen atom, (a) one of the two substituents of which Y is composed is not an ethyl group if the other substituent is a hydrogen atom, and (b) the group NY does not represent a morpholino or	15
20	piperidino heterocyclic ring; together with an appropriate carrier and at least one oxidation base. 6. A composition according to claim 5 which contains an oxidising agent at the time of use. 7. A composition according to claim 5 or 6, wherein the compound of the formula (I) is present in	20
_•	the composition in an amount of from 0.001 to 2.5% by weight, relative to the total weight of the composition. 8. A composition according to any one of claims 1 to 7 which contains at least one oxidation base	
	selected from para-phenylenediamines, para-aminophenols or heterocyclic bases. 9. A composition according to any one of claims 1 to 8 which contains at least one coupler	25
25	selected from resorcinol, pyrocatechol, 2-methylresorcinol, 2-ethylresorcinol, meta-aminophenol, 2-methyl-5-aminophenol, 2-methyl-5-N-(β-hydroxyethyl)-aminophenol, 6-hydroxybenzomorpholine, 2,6-dimethyl-3-acetylaminophenol, 2-methyl-5-carbethoxyaminophenol, 2-methoxy-5-carbethoxyamino-	
30	phenol, 2-methyl-5-ureidophenol, 2,4-diaminophenoxyethanol, 2,4-diaminoanisole, 2,6-dimethylmeta-phenylenediamine, 2-amino-4-N-methylaminophenoxyethanol, 2,4-diaminophenyl β -methoxyethyl ether, 2,4-diaminophenyl β -mesylaminoethyl ether, 2-N-carbamylmethylamino-4-amino-anisole, 3-amino-4-methoxyphenol, α -naphthol, 2,6-diaminopyridine, 3,5-diamino-2,6-	30
	dimethoxypyridine, 1-phenyl-3-methylpyrazol-5-one and 2-amino-4-N-(β -hydroxyethyl)-aminophenyl β -hydroxypropyl ether.	•
35	10. A composition according to any one of claims 1 to 9, which contains a compound selected from ortho-diphenol, ortho-phenylenediamines and ortho-aminophenols, the ortho-phenylenediamines and ortho-aminophenols being unsubstituted or substituted on the nucleus or on the amino group(s). 11. A composition according to any one of claims 1 to 10 which contains a dyestuff precursor of	35
40	the benzene series, containing, on the nucleus, at least three substituents selected from hydroxyl, methoxy and amino. 12. A composition according to any one of claims 1 to 11 which contains a quinone dyestuff.	40
	13. A composition according to any one of claims 1 to 12, which contains an indoaniline, indophenol, indamine or a leuco derivative thereof. 14. A composition according to any one of claims 1 to 13, which contains a dyestuff selected	•
45	from azo dyestuffs, anthraquinone dyestuffs and nitro dyestuffs of the benzene series. 15. A composition according to any of claims 1 to 14, which has a pH of from 5 to 11.5. 16. A composition according to any one of claims 1 to 15, which contains at least one adjuvant	45
50	selected from penetrating agents, surface-active agents, thickeners, antioxidants, alkalising or acidifying agents, perfumes, sequestering agents, film-forming products, treating agents and solvents. 17. A composition according to any one of claims 1 to 16, which is in the form of a liquid, a cream, a gel or an aerosol.	50
	18. A dyeing composition substantially as hereinbefore described in any one of Examples 12 to 41. 19. A hair-dyeing process comprising leaving a composition as claimed in claim 1 to act on the	
55	and the state of t	55
	oxidation base and at the time of use, a sufficient amount of oxidising agent is added to the dyeing composition used.	
60	21. A hair-dyeing process substantially as her inb for described in any one of Examples 12 to	60